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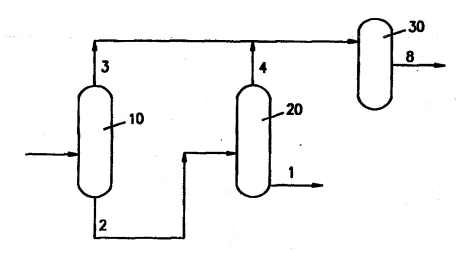
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(71) Applicant: EXXON RESEARCH AND ENGINEERING COMPANY [US/US]; P.O. Box 390, 180 Park Avenue, Florham Park, NJ 07932-0390 (US).

(72) Inventors: COOK, Bruce, Randall; 2 Capoolong Creek Road, Pittstown, NJ 08867 (US). BERLOWITZ, Paul, Joseph; 939 Jamestown Road, East Windsor, NJ 08520 (US). WITTENBRINK, Robert, Jay; 836 Shadyglen Drive, Baton Rouge, LA 70816 (US).

(74) Agents: SIMON, Jay et al.; Exxon Research and Engineering Company, P.O. Box 390, Florham Park, NJ 07932-0390

(54) Title: DIESEL ADDITIVE FOR IMPROVING CETANE, LUBRICITY, AND STABILITY



#### (57) Abstract

A process for producing additive compositions, especially via a Fischer-Tropsch reaction, useful for improving the cetane number or lubricity, or both the cetane number and lubricity, of a mid-distillate, diesel fuel. In producing the additive, the product of a Fischer-Tropsch reaction is separated into a high boiling fraction and a low boiling, e.g., a 700 °F-fraction. The high boiling fraction is hydroisomerized at conditions sufficient to convert it to a 700 °F-low boiling fraction, the latter being blended with the 700 °F-fraction and the diesel additive is recovered therefrom.

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# DIESEL ADDITIVE FOR IMPROVING CETANE, LUBRICITY, AND STABILITY

#### FIELD OF THE INVENTION

This invention relates to an additive for diesel fuels. More particularly, this invention relates to an additive that can provide cetane improvement, lubricity improvement and stability of diesel fuels regardless of their hydrocarbon source, i.e., natural or synthetic crudes.

#### BACKGROUND OF THE INVENTION

The continuing pressure from regulatory agencies around the world for reducing emissions, e.g., particulates, from diesel engines has lead to increased demand for high cetane diesel fuels. This demand has been met, but only in part, by blending refinery streams, e.g., raw or hydrotreated cat cracker, coker distillate, and virgin distillates that contain few, if any, paraffins with distressed streams of low native cetane. Also, cetane of refinery streams can be improved with severe hydrotreating which is expensive and limits cetane to the mid-fifties. Alternatively, commercial cetane additives, e.g., alkyl nitrates and peroxides, are available but expensive, often toxic, and therefore, limited as to the amount that can be used. Consequently, there is a need for an environmentally benign material that can significantly increase cetane, for example increasing cetane number leads to decreasing emissions of pollutants. Further, in severely hydrotreated materials lubricity is often inadequate and lubricity additives are required, too.

#### SUMMARY OF THE INVENTION

In accordance with this invention a diesel fuel additive that contributes cetane, lubricity, and stability to diesel fuel blends can be prepared from the Fischer-Tropsch hydrocarbon synthesis process, preferably a non-shifting process.

The diesel additive which can be blended with diesel fuel streams in amounts of at least about 1wt% can be described as

- boiling range 540-680°F;
- $\ge 90$  wt%  $C_{16}$ - $C_{20}$  paraffins, of which greater than 50 wt% are isoparaffins having substantial, i.e.,  $\ge 25$  wt%, mono-methyl paraffins;
  - cetane number of  $\geq$  87;
  - ≥ 2500 ppm as oxygen of  $C_{14}$ - $C_{16}$  linear, primary alcohols.

Additionally, such materials contain few unsaturates, e.g.,  $\leq 1$  wt% ppm total unsaturates (olefins + aromatics), preferably less than about 0.5 wt%; and nil sulfur and nitrogen, e.g.,  $\leq 50$  ppm by wt S or N. These materials are readily produced via a non-shifting Fischer-Tropsch (F/T) catalytic process followed by hydroisomerizing at least a portion of the heavier portion of the F/T product and blending it back with at least a portion of a lighter non-isomerized fraction and recovering the desired material.

#### BRIEF DESCRIPTION OF THE DRAWING

Figure 1 is a schematic representation of a process for producing the desired diesel fuel additive.

The diesel material of this invention, preferably produced in accordance with the process described herein, is best employed as a blending agent with other diesel fuels in need of upgrading, that is, upgrading or increasing cetane number, increasing lubricity, increasing stability, or any combination of the foregoing. The amount of additive employed will be that amount sufficient to improve the cetane or lubricity or both of the blend to meet desired specifications.

More preferably, diesel materials having a cetane number in the range 30-55, preferably less than about 50, preferably less than about 40 or diesel materials having lubricity measurements of less than 2500 grams in the scuffing BOCLE test or greater than 450 microns wear scar in the High Frequency Reciprocating Rig (HFRR) test, or both low cetane and poor lubricity are excellent candidates for upgrading with the diesel fuel additive of this invention.

There is essentially no upper limit on the amount of additive that can be used other than economic limits. In general, the diesel additive of this invention is used as a blend with diesel materials that are or can be used as diesel fuels in amounts of at least about 1 wt%, preferably in amounts of about 1-50%, more preferably in amounts of about 2 to 30%, and still more preferably in amounts of about 5-20%. (For rough estimation purposes about 1% additive will increase cetane number by about 0.5; and about 2-10% additive will improve lubricity by about 20% in the scuffing BOCLE test.)

Examples of distressed diesel materials requiring upgrading are raw and hydrotreated cat cracker and coker distillates. These materials are usually low in cetane number, being less than about 50, sometimes less than about 40. Additionally, hydrotreated distillates in the diesel boiling range,

particularly where sultur and nitrogen are less than 50 wppm and oxygenates are nil, can have their lubricity increased by virtue of blending with the diesel additive of this invention.

The BOCLE test is described in Lacy, P.I. "The U.S. Army Scuffing Load Wear Test", Jan. 1, 1994 which is based in ASTM D5001.

The HFRR test is described in "Determination of Lubricity of Diesel Fuel by High Frequency Reciprocating Rig (HFRR) Test". ISO Provisional Standard, TC22/SC7N595, 1995 and in "Pending ASTM Method: Standard Test Method for Evaluating Lubricity of Diesel Fuels by the High-Frequency Reciprocating Rig (HFRR)" 1996.

This invention, as described in the embodiment shown in Figure 1 is based, in part, on the discovery that a fractionated, hydroisomerized product obtained from a non-shifting Fischer-Tropsch process does not behave in a usual fashion. That is, usually, as molecular weight increases, cetane number also increases. However, as the boiling point of a particular fraction increases after hydroisomerizing, the iso-to normal ratio also increases and as the iso/ normal ratio increases, the cetane number decreases. Consequently, with increasing molecular weight and increasing iso/ normal ratio, a maximum cetane number occurs for a particular fraction. Also, at this maximum cetane, the cloud point, which also increases with increasing molecular weight, is acceptable and that fraction contains virtually nil unsaturates (for stability) and linear, primary alcohols which impart lubricity.

In the practice of this invention, the paraffinic stream from the F/T reactor is split, or divided, into (i) a high boiling liquid fraction and (ii) a low boiling liquid fraction, the split being made nominally at temperature ranging

between about 675°F and about 725°F, preferably at about 700°F to produce a nominally 700°F+ liquid fraction and a 700°F- liquid fraction. The high boiling or preferred 700°F+ fraction (i) is mildly hydroisomerized and hydrocracked to produce a 700°F- boiling product which is then combined with the native low boiling, or 700°F- boiling liquid fraction (ii), and this mixture is then separated, i.e., suitably fractionated, to produce very stable, environmentally benign, nontoxic, mid-distillate, diesel fuel additive.

Referring to the Figure there is shown a schematic for producing the desired fraction that is useful as a diesel fuel improver. Hydrogen and carbon monoxide is fed in line 1 into Fischer-Tropsch reactor 10 at reaction conditions. From the reactor 10 a product is recovered and may, for example, be recovered as a lighter stream or a heavier stream. The split may be at nominally 250°F, preferably 500°F, more preferably 700°F. Consequently, in the most preferred embodiment the lighter stream may be a 700°F- while the heavier stream is a 700°F+, lines 3 and 2, respectively. The heavier stream is then hydroisomerized in reactor 20 from which a 700°F- stream is recovered in line 4 and combined with the lighter product of line 3. The combined stream is fractionated in fractionator 30 from which the desired diesel blending fraction is recovered in line 8. Additional 700°F+ material from line 6 can be recovered, and if desired, recycled to reactor 20 for the production of additional 700°F- material.

Non-shift F/T reaction conditions are well known to those skilled in the art and can be characterized by conditions that minimize the formation of carbon dioxide byproducts. Non-shift F/T conditions can be achieved by a variety of methods, including one or more of the following: operating at relatively low carbon monoxide partial pressures, that is, operating at hydrogen carbon monoxide ratios of at least about 1.7:1, preferably about 1.7:1 to about

2.5:1, more preferably at least about 1.9:1, and in the range 1.9:1 to about 2.3:1 with an alpha of at least about 0.88, preferably at least about 0.91; temperatures of about 175-400°C, preferably about 180-300°C; using catalysts comprising cobalt or ruthenium as the primary F/T catalysts, preferably supported cobalt or supported ruthenium, most preferably supported cobalt where the support may be silica, alumina, silica-alumina or Group IVB metal oxides, e.g., titania. Promoters may also be employed, e.g., rhenium, titanium, zirconium, hafnium.

Whereas various catalysts can be used to convert syngas to F/T liquids, supported cobalt and ruthenium catalysts are preferred in that they tend to produce primarily paraffinic products; especially cobalt catalysts which tend toward making a heavier product slate, i.e., a product containing  $C_{20}+$ . The product withdrawn from the F/T reactor is characterized as a waxy Fischer-Tropsch product, a product which contains  $C_5+$  materials, preferably  $C_{20}+$  materials, a substantial portion of which are normal paraffins. A typical product slate is shown in Table A and can vary by about  $\pm 10\%$  for each fraction.

TABLE A

Typical product slate from F/T process liquids:

		<u>Wt.%</u>
IBP-320°F		13
320-500°F		23
500-700°F	- ()	19
700-1050°F		34
1050°F+		<u>11</u>

Table B below lists some typical and preferred conditions for conducting the hydroisomerization reaction.

#### **TABLE B**

	TYPICAL	PREFERRED
CONDITION		
CONDITION	RANGE	RANGE
Temperature, °F	300-800	600-750
Pressure, psig	0-2500	500-1200
Hydrogen treat rate, SCF/B	500-5000	2000-4000
Hydrogen consumption rate,		
SCF/B	50-500	100-300

While virtually any bifunctional catalyst may be satisfactorily used for conducting the hydroisomerization reaction, some catalysts perform better than others and are preferred. For example, catalysts containing a supported Group VIII non-noble metal, e.g., platinum or palladium, are useful as are catalysts containing one or more Group VIII metals, e.g., nickel, cobalt, which may or may not also include a Group VI metal, e.g., molybdenum. Group IB metals can also be used. The support for the metals can be any acidic oxide or zeolite or mixtures thereof. Preferred supports include silica, alumina, titania, zirconia, vanadia and other Group III, IV, VA or VI oxides, as well as Y sieves, such as ultrastable Y sieves. Preferred supports include alumina and silica-alumina. More preferred catalysts and supports are those described in U.S. Pat. No. 5,187,138 incorporated herein by reference. Briefly, the catalysts described therein contain one or more Group VIII metals on alumina or silica-alumina supports where the surface of the support is modified by addition of a silica

precursor, e.g., Si(OC<sub>2</sub>H<sub>5</sub>)<sub>4</sub>. Silica addition is at least 0.5 wt.% preferably at least 2 wt.%, more preferably about 2-25%.

In hydroisomerization reactions increasing conversion tends to increase cracking with resultant higher yields of gases and lower yields of distillate fuels. Consequently, conversion is usually maintained at about 35-80% of 700°F+ feed hydrocarbons converted to 700°F- hydrocarbons.

In one aspect, the 700°F- paraffinic mixture obtained from the F/T reactor is fractionated to produce an environmentally friendly, benign, non-toxic additive boiling within the range of from about 540°F to about 680°F, preferably from about 570°F to about 650°F, which when combined with mid-distillate, diesel fuels will produce products of outstanding lubricity. These additives will contain generally more than 90 wt%, preferably more than 95 wt%, and more preferably more than 98 wt%, C<sub>16</sub> to C<sub>20</sub> paraffins, based on the total weight of the additive, of which greater than 50 wt%, based on the total weight of the paraffins in the mixture, are isoparaffins; and the isoparaffins of the mixture are further defined as greater than 25 percent, preferably greater than 40 percent, and more preferably greater than 50 percent, by weight, mono-methyl paraffins. The additive composition is also rich in  $C_{14}$ - $C_{16}$  linear primary alcohols species which impart higher lubricity, when combined with a mid-distillate, diesel fuel. In general the linear primary alcohols constitute at least about 0.05 percent, preferably at least about 0.25 percent, and generally from about 0.25 percent to about 2 percent, or more, of the additive mixture, based on the total weight of the additive.

#### Example 1

- a) A mixture of hydrogen and carbon monoxide synthesis gas (H<sub>2</sub>:CO 2.11-2.16) was converted to heavy paraffins in a slurry Fischer-Tropsch reactor. A titania supported cobalt/rhenium catalyst was utilized for the Fischer-Tropsch reaction. The reaction was conducted at 422-428°F, 287-289 psig, and the feed was introduced at linear velocity of 12 to 17.5 cm/sec. The alpha of the Fischer-Tropsch synthesis step was 0.92. The paraffinic Fischer-Tropsch product was isolated in three nominally different boiling streams, separated by utilizing a rough flash. The three boiling fractions obtained were: 1) a native low boiling C<sub>3</sub>-500°F fraction, i.e., F/T cold separator liquids; 2) a 500-700°F boiling fraction, i.e., F/T hot separator liquids, and 3) a 700°F+ boiling fraction, i.e., or F/T reactor wax.
- b) The 700°F+ boiling fraction, or F/T reactor wax, having a boiling point distribution as follows: IBP-500°F, 1.0%, 500°F-700°F, 28.1%, and 700°F+, 70.9%, was then hydroisomerized and hydrocracked over a dual functional catalyst consisting of cobalt (CoO, 3.2 wt.%) and molybdenum (MoO<sub>3</sub>, 15.2 wt.%) on a silica-alumina cogel acidic support, 15.5 wt.% of which is SiO<sub>2</sub> to obtain a 700°F- product. The catalyst had a surface area of 266 m/g and pore volume (PV<sub>H2O</sub>) of 0.64 ml/g. The conditions for the reaction are listed in Table 1A and were sufficient to provide approximately 50% 700°F+ conversion where 700°F+ conversion is defined as 700°F+ Conv.= [1 (wt.% 700°F+ in product)/(wt.% 700°F+ in feed)] x 100

- 10 -

#### TABLE 1A

## **Operating Conditions**

Temp., °F

690

LHSV, v/v/h

0.6 - 0.7

H<sub>2</sub> Pressure, psig (pure)

725 2500

H<sub>2</sub> Treat rate, SCF/B

- c) To simulate the total of the 700°F- liquids derived in steps (a) and (b), above, seventy-eight wt.% hydroisomerized F/T reactor wax boiling at 700°F-, 12 wt.% F/T cold separator liquids, and 10 wt.% F/T hot separator liquids from a large scale pilot unit were combined and mixed. A final diesel fuel, i.e., a 250-700°F boiling fraction was isolated by distillation from this blend. The hydroisomerized F/T reactor wax was prepared in flow through, fixed bed unit using a cobalt and molybdenum promoted amorphous silicalumina catalyst, as described in U.S. Patent 5,292,989 and U.S. Patent 5,378,348.
- d) The diesel fuel of step (c), above, was fractionated using a 15/5 distillation column into 9 cuts of increasing boiling range. These cuts, the mid-boiling points and engine cetane number of each fraction are listed in Table 1B. A composite 33%-55% volume fraction was also made and is shown in this table.

TABLE 1B

Cut#	Volume Fraction	Initial B.P. (°F)	50% B.P.	Final B.P.	Engine Cetane Number
1	0-10%	206	317	383	60.7
2	10-20%	294	398	469	70.5
3	20-30%	354	461	536	77.4
4	30-40%	419	515	560	83.2
5	40-50%	461	551	590	84.3
6	50-60%	494	578	612	84.1
7	60-70%	544	610	645	88.5
8	70-80%	571	641	676	87.9
9	80- 100%	605	691	737	81.6
	33-55%	500		570	84
	60-80%	570	•	670	88

All of the fractions, as clearly evident, exhibit high engine cetane numbers, with fractions 7 and 8 having the highest cetane. The cetane number of a composite of the 33-55% volume fraction has a cetane number of 84. Cetane number is clearly not simply a function of boiling point, as the highest boiling fraction 9 has a significantly lower cetane number than 7 and 8. The 33-55% composite fraction, and 60-80% composite fractions were in fact found to

contain distinctive molecular compositions that lead to these improved properties.

In Table 1C is given a projected combination of Fractions 7 + 8 (60%-80%), from the analysis of the individual fractions by GC and GC/MS. The linear primary alcohol content leads to improved lubricity; lubricity increasing as the alcohol content of the fraction is increased.

TABLE 1C

1.34

Wt.% Paraffin Carbon		
C <sub>15</sub>	0.2	
C <sub>16</sub>	3.2	
C <sub>17</sub>	22.4	
C <sub>18</sub>	37.5	
C <sub>19</sub>	28.4	
C <sub>20</sub>	8.0	

 $C_{21}$ 0.2

wppm linear primary alcohols:

Iso/Normal

 $C_{14}$ 267 C<sub>15</sub> 1740  $C_{16}$ 1024

In Table 1D is given a projected combination of cuts 4, 5 and 6 which encompasses the 33-55% volume fraction. Analysis of the individual fractions by GC and GC/MS show that the fractions contain relatively high concentrations of linear primary alcohols. The linear primary alcohol content leads to improved lubricity; lubricity increasing as the alcohol content of the fraction is increased.

## TABLE 1D

Wt.% Paraffin Car	bon
C <sub>14</sub>	2.8
$C_{16}$	54.8
C <sub>17</sub>	42.3
Iso/Normal	1.21
wppm linear primary alcohols:	
C <sub>12</sub>	379
C <sub>13</sub>	4404
C <sub>14</sub>	1279

The following Table 1E is a further tabulation of tests performed on the 9 cuts, and a composite of the 9 cuts, showing the lubricity in terms of the BOCLE test, the Peroxide No., and the cloud and pour points.

TABLE 1E

Cut	Lubricity <sup>1</sup>	Peroxide No. <sup>2</sup>	Cloud <sup>3</sup>	Pour <sup>4</sup>
1	33	76.0 (Fail)	<-49	<-49
2	35	6.7 (Fail)	<-45	<-45
3	55	2.0 (Fail)	<-27	<-28
4	73	0.6 (Pass)	<-15	<-15
5	75	0.9 (Pass)	-4	-3
6	93	0.7 (Pass)	2	3
7	102	0.3 (Pass)	6	6
8	117	0.0 (Pass)	8	9
9	129	0.4 (Pass)	13	12
Sum Cuts 1-9 <sup>5</sup>	75	7.5 (Pass)	-8	-8
33-55% Volume Fraction <sup>6</sup>	>75	<1 (Pass)	<-5	<-5

#### Notes:

- Lubricity results in the BOCLE test as described in Lacy, P.I. "The U.S. Army Scuffing Load Wear Test", Jan. 1, 1994 which is based in ASTM D5001. Results are represented as a % of the high reference fuel, Cat 1-K specified in the procedure.
- 2 Peroxide number according to ASTM D3703. 100 mls of fuel were filtered, then aerated for 3 minutes with air, and then placed in a brown 4 oz. bottle in a 65C oven for 4 weeks. Peroxide number was measured at the start of

the test, and after 7, 14, 21 and 28 days. At the end of the test those fuels with peroxide number <1 were considered to have good stability and passed the test.

- 3 Cloud point as described by ASTM D2500.
- 4 Pour point as described by ASTM D97.
- 5 Entire product of cuts 1 through 9 before fractionation.
- 6 Estimation from result from cuts 4-6, as a neat fuel.

These data thus show materials which can provide significant benefits to cetane number and lubricity without incurring debits due to oxidative instability or excessively high cloud/pour points. Blending this additive into a base 35 cetane stream at 5-10% produces cetane number improvements of 2.5 to 5 numbers with improved lubricity and essentially no effect on cold flow properties.

#### CLAIMS:

- 1. A diesel fuel additive comprising
- (i)  $\geq$  90 wt% C<sub>16</sub>-C<sub>20</sub> paraffins, of which  $\geq$  50% are isoparaffins at least a portion of which are mono-methyl branched;
  - (ii) cetane number of ≥ 87;
  - (iii)  $\geq$  2500 ppm as oxygen of C<sub>14</sub>-C<sub>16</sub> linear, primary alcohols;
  - (iv) a boiling range of 540-680°F.
- 2. The additive of claim 1 wherein the paraffins are  $\geq$  95 wt%, and the mono-methyl branched isoparaffins are  $\geq$  25 wt%.
- 3. The additive of claim 2 wherein the  $C_{14}$ - $C_{16}$  alcohols are present in an amount of 0.25 to 2 wt%.
- The additive of claim 2 wherein the sulfur and nitrogen concentrations are each ≤ 50 wppm and the unsaturates concentration ≤ 1 wt%.
- 5. The additive of claim 1 derived from a non-shifting Fischer-Tropsch process.
- 6. The additive of claim 1 combined with material in an amount of 1-50 wt%.
- 7. The combination of claim 6 wherein the diesel material has a cetane of  $\leq 50$ .

- 8. The combination of claim 6 wherein the diesel material has a lubricity of less than 2500 grams in the scuffing BOCLE test.
- 9. The combination of claim 6 wherein the additive is present in an amount of about 2-30 wt%.
- 10. The combination of claim 6 wherein the diesel material is selected from the group consisting of raw and hydrotreated cat cracker and coker distillates having a cetane number  $\leq$  40 and hydrotreated distillates in the diesel boiling range having a lubricity of less than 2500 grams in the scuffing BOCLE test.
- 11. A process for preparing a diesel fuel additive described in claim 1 comprising
- (a) reacting hydrogen and carbon monoxide at reaction conditions in the presence of a non-shifting Fischer-Tropsch catalyst,
- (b) recovering at least a portion of the liquid product of the reaction and separating at least a portion of the liquid product into a heavier fraction and a lighter fraction,
- (c) hydroisomerizing at hydroisomerization conditions at least a portion of the heavier fraction and recovering a 700°F- product,
- (d) combining the lighter fraction of step (b) with the 700°F-product of step (c) and recovering a diesel fuel additive.
- 12. The process of claim 1 wherein the heavier fraction of step (b) is a 675°F+ material.

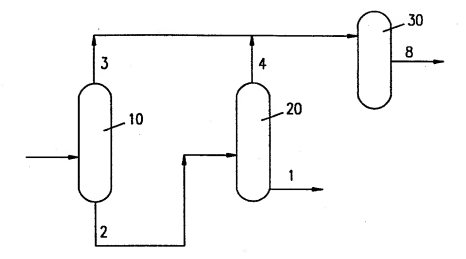


FIG. 1

# INTERNATIONAL SEARCH REPORT

national Application No PCT/US 98/01670

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A. CLASSI IPC 6	FICATION OF SUBJECT MATTER C10L1/08 C10L1/18 C10L1C	)/04	
According to	o International Patent Classification(IPC) or to both national class	ification and IPC	
	SEARCHED		
Minimum do IPC 6	cumentation searched (classification system followed by classific C10L C10G	cation symbols)	, .
Documenta	tion searched other than minimumdocumentation to the extent th	at such documents are included in the fields se	arched
Electronic d	ata base consulted during the international search (name of data	base and, where practical, search terms used	)
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.
Ρ,Χ	WO 97 14769 A (EXXON) 24 April see claims 5,12	1997	11
A	EP 0 321 301 A (EXXON) 21 June see examples 1-3	1989	11,12
A	EP 0 587 246 A (SHELL) 16 March see column 7	1994	11
A	FR 859 686 A (SYNTHETIC OILS LT 24 December 1940 see the whole document	TD ET AL.)	11
A	FR 732 964 A (DEUTSCHE HYDRIERW 28 September 1932 see the whole document	WERKE AKT.)	1
		-/ <b></b>	
X Furt	her documents are listed in the continuation of box C.	X Patent family members are listed	in annex.
"A" docum consider "E" earlier filling of "L" docum which citatio "O" docum other "P" docum later t	ent which may throw doubts on priority claim(s) or is cited to establish the publicationdate of another in or other special reason (as specified) sent referring to an oral disclosure, use, exhibition or means ent published prior to the international filing date but than the priority date claimed	"T" later document published after the inte or priority date and not in conflict with cited to understand the principle or th invention  "X" document of particular relevance; the cannot be considered novel or canno involve an inventive step when the de "Y" document of particular relevance; the cannot be considered to involve an in document is combined with one or ments, such combination being obvious in the art.  "&" document member of the same patent	the application but secry underlying the claimed invention to considered to coursent is taken alone claimed invention eventive step when the ore other such docurus to a person skilled
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Jalegury	опшного осошнени, или вымошностичете аругориала, от ша тогачали разбадав		Constant Grant IIV.
Ą	WO 96 26994 A (EXXON) 6 September 1996 see the whole document		. 1
A	FR 2 650 289 A (LASCARAY,S.A.) 1 February 1991 see the whole document		1
<b>A</b>	US 5 324 335 A (BENHAM ET AL.) 28 June 1994	***	
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#### INTERNATIONAL SEARCH REPORT

Information on patent family members

ational Application No PCT/US 98/01670

Patent document cited in search repor	<b>1</b>	Publication date	Patent family member(s)	Publication date
WO 9714769	A	24-04-1997	AU 7395196 A	07-05-1997
EP 321301	Α	21-06-1989	AU 2694488 A DE 3873291 A JP 1297148 A	22-06-1989 03-09-1992 30-11-1989
			US 4923841 A	08-05-1990
EP 587246	A	16-03-1994	AU 662437 B AU 4617993 A CA 2104913 A DE 69312607 D DE 69312607 T ES 2105082 T JP 6198180 A NO 933184 A ZA 9306543 A	31-08-1995 17-03-1994 09-03-1994 04-09-1997 15-01-1998 16-10-1997 19-07-1994 09-03-1994 28-03-1994
FR 859686	Α	26-12-1940	NONE	
FR 732964	Α	28-09-1932	NONE	
WO 9626994	Α	06-09-1996	CA 2213656 A EP 0812346 A	06-09-1996 17-12-1997
FR 2650289	Α	01-02-1991	NONE	
US 5324335	A	28-06-1994	WO 9520558 A AU 6028894 A US 5504118 A US 5645613 A US 5621155 A US 5506272 A US 5543437 A US 5500449 A US 5620670 A	03-08-1995 15-08-1995 02-04-1996 08-07-1997 15-04-1997 09-04-1996 06-08-1996 19-03-1996 15-04-1997